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=> s ((mrna or rna or vrna) (N) ((adna or cdna or dna)) (3n) hybrid? UNMATCHED LEFT PARENTHESIS '((MRNA' The number of right parentheses in a query must be equal to the number of left parentheses.

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 6 FILES SEARCHED...
- L1 31952 ((MRNA OR RNA OR VRNA) (N) (ADNA OR CDNA OR DNA)) (3N) HYBRID?
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- L2 10362864 L1 (S) INHIB? OR REDUC?
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- L3 3102 L1 (S) (INHIB? OR REDUC?)
- => s l1 (s) ((inhib? or reduc?) (n) expres?)
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- 5 FILES SEARCHED...
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L5 67 DUP REM L4 (0 DUPLICATES REMOVED)

=> s 15 <=2001

NUMERIC EXPRESSION NOT VALID 'L34 <=2001'

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Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

(REVIEW, TUTORIAL)

LANGUAGE: English

DOCUMENT TYPE:

FILE SEGMENT: Priority Journals

ENTRY MONTH: 200210

ENTRY DATE: Entered STN: 20020822

Last Updated on STN: 20021019 Entered Medline: 20021001

AB D-RNAi (Messenger RNA-antisense DNA interference), a novel posttranscriptional phenomenon of silencing gene expression by

transfection of mRNA-aDNA hybrids, was

originally observed in the effects of bcl-2 on phorbol ester-induced apoptosis in human prostate cancer LNCaP cells. This phenomenon was also demonstrated in chicken embryos and a human CD4(+) T cell line, H9. The in vivo transduction of beta-catenin D-RNAi was shown to knock out more than 99% endogenous beta-catenin gene expression, while the in cell transfection of HIV-1 D-RNAi homolog rejected viral gene replication completely. D-RNAi was found to have long-term gene knockout effects resulting from a posttranscriptional gene silencing mechanism that may involve the homologous recombination between intracellular mRNA and the mRNA components of a D-RNAi construct. These findings provide a potential intracellular defense system against cancer and viral infections.

L15 ANSWER 2 OF 21 MEDLINE ON STN ACCESSION NUMBER: 2001216119 MEDLINE DOCUMENT NUMBER: PubMed ID: 11237705

TITLE: A Novel mRNA-cDNA interference phenomenon for silencing

bcl-2 expression in human LNCaP cells.

AUTHOR: Lin S L; Chuong C M; Ying S Y

CORPORATE SOURCE: Department of Pathology, Keck School of Medicine,

University of Southern California, HMR-209, 2011 Zonal

Avenue, Los Angeles, California, 90033, USA.

SOURCE: Biochemical and biophysical research communications,

(2007 18-10) 201 (2) (20 44

(2001 Mar 2) 281 (3) 639-44.

Journal code: 0372516. ISSN: 0006-291X.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200104

ENTRY DATE:

Entered STN: 20010425

Last Updated on STN: 20010425 Entered Medline: 20010419

The templates required for inducing posttranscriptional gene silencing (PTGS) effects have been investigated in human prostate cancer LNCaP cells. Transfection of a mRNA-cDNA hybrid construct was found to result in a relatively long-term interference of specific gene expression. Androgen-stimulated expression of bcl-2 has been reported to increase the tumorigenic and metastatic potentials of human prostate cancer LNCaP cells, as well as their resistance to many apoptotic stimuli. The addition of bcl-2 antisense oligonucleotides, however, restored apoptosis. Our studies demonstrate gene silencing effects of the mRNA-cDNA transfection that is similar to those of PTGS/RNAi in this in vitro prostate cancer cell model. A potential RNA-directed RNA polymerase activity was also detected which is alpha-amanitin-sensitive. These findings indicate that a novel gene

silencing system may exist in mammalian cells. Copyright 2001 Academic Press.

L15 ANSWER 3 OF 21 BIOSIS COPYRIGHT (c) 2005 The Thomson Corporation on STN

ACCESSION NUMBER: 1999:78994 BIOSIS DOCUMENT NUMBER: PREV199900078994

TITLE: Lipid A mutant Salmonella with suppressed virulence and

TNFalpha induction retain tumor-targeting in vivo.

AUTHOR(S): Low, K. Brooks; Ittensohn, Martina; Le, Trung; Platt, James; Sodi, Stefano; Amoss, Max; Ash, Olivia; Carmichael,

Ellen; Chakraborty, Ashok; Fischer, Jessica; Lin, Stanley L.; Luo, Xiang; Miller, Samuel I.; Zheng, Li-Mou; King, Ivan; Pawelek, John M.; Bermudes, David

[Reprint author]

CORPORATE SOURCE: Vion Pharm. Inc., New Haven, CT 06511, USA

SOURCE: Nature Biotechnology, (Jan., 1999) Vol. 17, No. 1, pp.

37-41. print. ISSN: 1087-0156.

DOCUMENT TYPE: Article LANGUAGE: English

ENTRY DATE: Entered STN: 1 Mar 1999

Last Updated on STN: 1 Mar 1999

AΒ Systemically administered tumor-targeted Salmonella has been developed as an anticancer agent, although its use could be limited by the potential induction of tumor necrosis factor a (TNFalpha)-mediated septic shock stimulated by lipid A. Genetic modifications of tumor-targeting Salmonella that after lipid A and increase safety must, however, retain the useful properties of this bacteria. We report here that disruption of the Salmonella msbB gone reduces TNFalpha induction and increases the LD50 of this pathogenic bacteria by 10,000-fold. Notwithstanding this enormous difference, Salmonella retains its tumor-targeting properties, exhibiting tumor accumulation ratios in excess of 1000: 1 compared with normal tissues. Administration of this bacteria to mice bearing melanoma results in tumors that are less than 6% the size of tumors in untreated controls at day 18. Thus, the antitumor activity previously demonstrated using tumor-targeting Salmonella with normal lipid A is retained. Lipid modification of tumor-specific bacterial vectors provides a means for reducing septic shock and further suggests that the antitumor activity of these bacteria may be independent of TNFalpha.

L15 ANSWER 4 OF 21 SCISEARCH COPYRIGHT (c) 2005 The Thomson Corporation on

STN

ACCESSION NUMBER: 2001:643563 SCISEARCH

THE GENUINE ARTICLE: 460VB

TITLE: Combined blockade of protein kinase A and bcl-2 by

antisense strategy induces apoptosis and inhibits tumor

growth and angiogenesis

AUTHOR: Tortora G (Reprint); Caputo R; Damiano V; Bianco R;

Fontanini G; Cuccato S; De Placido S; Bianco A R;

Ciardiello F

CORPORATE SOURCE: Univ Naples Federico II, Dipartimento Endocrinol & Oncol

Mol & Clin, Cattedra Oncol Med, Via S Pansini 5, I-80131

Naples, Italy (Reprint); Univ Naples Federico II,

Dipartimento Endocrinol & Oncol Mol & Clin, Cattedra Oncol

Med, I-80131 Naples, Italy; Univ Pisa, Ist Anat Patol,

Dipartimento Oncol, I-56100 Pisa, Italy

COUNTRY OF AUTHOR: Italy

SOURCE: CLINICAL CANCER RESEARCH, (AUG 2001) Vol. 7, No.

8, pp. 2537-2544.

Publisher: AMER ASSOC CANCER RESEARCH, PO BOX 11806,

BIRMINGHAM, AL 35202 USA.

ISSN: 1078-0432. Article; Journal

LANGUAGE: English

REFERENCE COUNT: 39

DOCUMENT TYPE:

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

AB Protein kinase A type I (PKAI) plays a key role in neoplastic transformation, conveys mitogenic signals from different sources, and is overexpressed in the majority of human tumors. Inhibition of PKAI by different tools results in cancer-cell growth inhibition in vitro and in vivo. We and others have recently shown that a novel class of mixed-backbone oligonucleotides targeting the PKAI subunit RI alpha exhibits improved pharmacokinetic properties and antitumor activity

accompanied by increased apoptosis in several human cancer types in vitro and in vivo. The role of bcl-2 in the control of apoptosis has been widely documented, and the inhibition of bcl-2 expression and function may have important therapeutic implications. In fact, oligonucleotides antisense bcl-2 have shown antitumor activity in animal models and have successfully completed early clinical trials. Recent studies have demonstrated a direct role of PKA in the regulation of the bcl-2-dependent apoptotic pathway. Therefore, we have investigated the combined blockade of PKA and bcl-2 by antisense strategy as a potential therapeutic approach. The novel hybrid DNA/RNA mixed-backbone oligonucleotide

antisense Riot (AS RI alpha) in combination with the antisense bcl-2 (AS bcl-2), cooperatively inhibited bcl-2 expression and soft agar growth and induced apoptosis in different human cancer cell lines. p.o. administration of AS RI alpha in combination with i.p. AS bcl-2 caused a marked antitumor effect and a significant prolongation of survival in nude mice bearing human colon cancer xenografts. Moreover, histochemical analysis of tumor specimens showed inhibition of RI alpha and Ki67 expression, inhibition of angiogenesis, and parallel induction of apoptosis in vivo. The results of our study imply an

induction of apoptosis in vivo. The results of our study imply an interaction between the PKA and bcl-2 signaling pathways and, because both antisenses have now entered Phase II trials, provide the rationale to translate this novel therapeutic strategy in a clinical setting.

L15 ANSWER 5 OF 21 EMBASE COPYRIGHT 2005 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 78062896 EMBASE

DOCUMENT NUMBER: 1

1978062896

TITLE:

Endonuclease V of Escherichia coli.

AUTHOR:

Gates III F.T.; Lin S.

CORPORATE SOURCE:

Dept. Biochem., Univ. California, Berkeley, Calif. 94720,

United States

SOURCE:

Journal of Biological Chemistry, (1977) Vol. 252, No. 5,

pp. 1647-1653. CODEN: JBCHA3

DOCUMENT TYPE:

Journal

FILE SEGMENT:

004 Microbiology

029 Clinical Biochemistry

LANGUAGE: English

AB A small endodeoxyribonuclease (2.3 S) that is active on single stranded DNA has been extensively purified from E. coli so as to be free of other known DNases. It has an alkaline pH optimum (9.5), requires Mg2+, and makes 3' hydroxy and 5' phosphate termini. The nuclease nicks duplex DNA, particularly if treated with OsO4, irradiated with ultraviolet light, or exposed to pH 5. The uracil containing duplex DNA from the Bacillus subtilis phage PBS 2 is an especially good substrate; it is made acid soluble by levels of the enzyme which fail to produce any acid soluble material in other single stranded or duplex DNAs. Neither RNA nor RNA DNA hybrid are degraded by the enzyme.

The enzyme specificity suggests that it might act at abnormal regions in DNA, so that its in vivo function could be to initiate an excision repair sequence. Its high activity on uracil containing DNA could imply that the enzyme provides an alternative mechanism for excising uracil residues from DNA to the pathway utilizing uracil DNA N glycosidase. The authors suggest that this enzyme be designated as endonuclease V of E. coli.

L15 ANSWER 6 OF 21 CA COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER:

136:145646 CA

TITLE:

Electronic detection of nucleic acids: A versatile

platform for molecular diagnostics

AUTHOR(S): Umek, Robert M.; Lin, Sharon W.; Vielmetter,

Jost; Terbrueggen, Robert H.; Irvine, Bruce; Yu, C. J.; Kayyem, Jon Faiz; Yowanto, Handy; Blackburn, Gary

F.; Farkas, Daniel H.; Chen, Yin-Peng

CORPORATE SOURCE:

Clinical Micro Sensors Division of Motorola, Inc.,

Pasadena, CA, 91105, USA

SOURCE:

Journal of Molecular Diagnostics (2001),

3(2), 74-84

CODEN: JMDIFP; ISSN: 1525-1578

PUBLISHER:

Association for Molecular Pathology

DOCUMENT TYPE:

Journal

LANGUAGE:

English

A novel platform for the electronic detection of nucleic acids on microarrays is introduced and shown to perform well as a selective detection system for applications in mol. diagnostics. A gold electrode in a printed circuit board is coated with a self-assembled monolayer (SAM) containing DNA capture probes. Unlabeled nucleic acid targets are immobilized on the surface of the SAM through sequence-specific hybridization with the DNA capture probe. A sep. signaling probe, containing ferrocene-modified nucleotides and complementary to the target in the region adjoining the capture probe binding site, is held in close proximity to the SAM in a sandwich complex. The SAM allows electron transfer between the immobilized ferrocenes and the gold, while insulating the electrode from soluble redox species, including unbound signaling probes. Here, we demonstrate sequence-specific detection of amplicons after simple dilution of the reaction product into hybridization buffer. In addition, single nucleotide polymorphism discrimination is shown. A genotyping chip for the C282Y single nucleotide polymorphism associated with hereditary hemochromatosis is used to confirm the genotype of six patients' DNA. In addition, a gene expression-monitoring chip is described that surveys five genes that are differentially regulated in the cellular apoptosis response. Finally, custom modification of individual electrodes through sequence-specific hybridization demonstrates the potential of this system for infectious disease diagnostics. The versatility of the electronic detection platform makes it suitable for multiple applications in diagnostics and pharmacogenetics.

REFERENCE COUNT:

16 THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L15 ANSWER 7 OF 21 USPATFULL on STN

ACCESSION NUMBER:

2001:93292 USPATFULL

TITLE:

Method for identifying essential or functional genes

INVENTOR(S):

Nilsen, Timothy W., Russell, OH, United States

PATENT ASSIGNEE(S):

Yale University, New Haven, CT, United States (U.S.

corporation)

NUMBER KIND DATE -----

PATENT INFORMATION:

US 6248525 B1 20010619 US 1998-196523 19981120 (9)

APPLICATION INFO.:

DOCUMENT TYPE:

Utility

FILE SEGMENT:

GRANTED Guzo, David

PRIMARY EXAMINER: ASSISTANT EXAMINER:

Leffers, Jr., Gerald G

LEGAL REPRESENTATIVE:

Arnall Golden Gregory LLP

NUMBER OF CLAIMS:

17

EXEMPLARY CLAIM: NUMBER OF DRAWINGS:

14 Drawing Figure(s); 11 Drawing Page(s)

LINE COUNT:

1527

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Two methodologies are provided: the first provides a means for rapidly and efficiently identifying essential and functional genes; and the second provides a means for obtaining biologically active nucleic molecules (ribozymes, EGSs, and antisense,) which can be used to inactivate functional genes. In the first method, a library of EGSs is prepared based on all possible known compositions. In a preferred embodiment, the EGSs are twelve or thirteen-mers for targeting bacterial RNAse to cleave a substrate. This library is added to the cells containing the genes to be screened, for example, E. coli. Those cells in which the EGS causes a loss of viability, or other phenotype, are identified. The EGS(s) responsible for the loss of viability are analyzed, and the resulting sequence information used to identify the gene within the known genomic sequences. In the second method, nucleotide molecules with optimal biological activity, for example, directing cleavage of a gene of interest by RNase P, are rapidly identified through the use of a vector including two reporter genes, the first in phase with the gene of interest, and the second as a control to verify that the vector is present in a cell or to aid in selection of cells containing the vector. Those cells where the gene of interest is cleaved by the functional oligonucleotide molecule can then be identified by reference to reporter gene 1. The responsible functional oligonucleotide molecules is then isolated and characterized.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 8 OF 21 USPATFULL on STN

ACCESSION NUMBER: 1999:7488 USPATFULL

TITLE:

Antisense oligonucleotides to inhibit expression of

mutated and wild type genes for collagen

INVENTOR(S): Prockop, Darwin, Philadelphia, PA, United States

Colige, Alain, Sart Tilman Par Liege, Belgium Baserga, Renato, Ardmore, PA, United States Nugent, Paul, Philadelphia, PA, United States

PATENT ASSIGNEE(S): Thomas Jefferson University, Philadelphia, PA, United

States (U.S. corporation)

	NUMBER	KIND DATE	
PATENT INFORMATION:	US 5861502	19990119	<
	WO 9411494	19940526	· <
APPLICATION INFO.:	US 1995-432158	19950630	(8)
	WO 1993-US10756	19931109	
		19950630	PCT 371 date
		19950630	PCT 102(e) date

RELATED APPLN. INFO.: Continuation of Ser. No. US 1992-973832, filed on 9 Nov

1992, now abandoned

DOCUMENT TYPE: Utility

FILE SEGMENT: Granted PRIMARY EXAMINER: Fox, David T.

ASSISTANT EXAMINER: Nelson, Amy J. LEGAL REPRESENTATIVE: Pennie & Edmonds LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 3 Drawing Figure(s); 3 Drawing Page(s)

LINE COUNT: 1486

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention is directed to oligonucleotides that inhibit mutant COL1A1 and/or wild type COL1A1 gene expression. The present invention is further directed to methods of inhibiting mutant and/or wild type collagen gene expression using the disclosed inhibitory oligonucleotides. The oligonucleotides and methods of the present invention are useful for the treatment of mammals having diseases related to inappropriate mutant or wild type COLIA1 gene expression.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 9 OF 21 USPATFULL on STN

ACCESSION NUMBER: 1998:68835 USPATFULL

TITLE: Expression of the developmental I antigen by a cloned

human cDNA encoding a member of a beta-1,

6-N-acetylglucosaminyltransfrase gene family INVENTOR(S):

Fukuda, Minoru, San Diego, CA, United States Bierhuizen, Marti F. A., Schiedam, Netherlands

PATENT ASSIGNEE(S): La Jolla Cancer Research Foundation, La Jolla, CA,

United States (U.S. corporation)

KIND NUMBER -----

PATENT INFORMATION: US 5766910

19980616 19950607 (8) US 1995-488135 APPLICATION INFO.:

RELATED APPLN. INFO.: Division of Ser. No. US 1993-118906, filed on 9 Sep

1993, now patented, Pat. No. US 5484590

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Prouty, Rebecca E. LEGAL REPRESENTATIVE: Campbell & Flores LLP

NUMBER OF CLAIMS: 10 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 18 Drawing Figure(s); 12 Drawing Page(s)

LINE COUNT: 1409

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides an isolated nucleic acid molecule encoding both a soluble and membrane-bound human β -1,6-Nacetylglucosaminyltransferase, the I-branching enzyme (IGnT). The invention also provides vectors containing the isolated nucleic acid molecule encoding human IGnT as well as recombinant host cells transformed with the vectors. The invention further provides a method of preparing a membrane-bound form of human IGnT and methods of preparing and purifying soluble human IGnT and active fragments of either form. Also provided are antisense oligonucleotides complementary to a nucleic acid molecule encoding a human IGnT or an active fragment thereof, antibodies directed to the human IGnT, pharmaceutical compositions related to the human IGnT and transgenic nonhuman mammals expressing DNA encoding normal or mutant human IGnT. Also provided are methods for regulating the expression of human IGnT and methods for modifying a biological function mediated by the regulatory activity of human IGnT. Methods of detecting the presence of linear polylactosaminoglycans

expressing i antigenic determinants on a cell surface also are provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 10 OF 21 USPATFULL on STN

ACCESSION NUMBER: 1998:31124 USPATFULL

TITLE: Antibodies to human I-branching beta-1,6-N-

acetylglucosaminyltransferase

INVENTOR(S): Fukuda, Minoru, San Diego, CA, United States

Bierhuizen, Marti F. A., Schiedam, Netherlands

PATENT ASSIGNEE(S): La Jolla Cancer Research Foundation, La Jolla, CA,

United States (U.S. corporation)

NUMBER KIND DATE

US 5731420 19980324 US 1995-486196 19950607 (8) PATENT INFORMATION:

APPLICATION INFO.:

Continuation of Ser. No. US 1993-118906, filed on 9 Sep RELATED APPLN. INFO.:

1993, now patented, Pat. No. US 5484590

DOCUMENT TYPE: Utility

FILE SEGMENT: Granted

PRIMARY EXAMINER: Cunningham, Thomas M. ASSISTANT EXAMINER: Lubet, Martha T. LEGAL REPRESENTATIVE: Campbell & Flores LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM: 1 NUMBER OF DRAWINGS:

15 Drawing Figure(s); 12 Drawing Page(s)

LINE COUNT:

1355

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

The present invention provides an isolated nucleic acid molecule encoding both a soluble and membrane-bound human β -1,6-Nacetylglucosaminyltransferase, the I-branching enzyme (IGnT). The invention also provides vectors containing the isolated nucleic acid molecule encoding human IGnT as well as recombinant host cells transformed with the vectors. The invention further provides a method of preparing a membrane-bound form of human IGnT and methods of preparing and purifying soluble human IGnT and active fragments of either form. Also provided are antisense oligonucleotides complementary to a nucleic acid molecule encoding a human IGnT or an active fragment thereof, antibodies directed to the human IGnT, pharmaceutical compositions related to the human IGnT and transgenic nonhuman mammals expressing DNA encoding normal or mutant human IGnT. Also provided are methods for regulating the expression of human IGnT and methods for modifying a biological function mediated by the regulatory activity of human IGnT. Methods of detecting the presence of linear polylactosaminoglycans expressing i antigenic determinants on a cell surface also are provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 11 OF 21 USPATFULL on STN

ACCESSION NUMBER:

96:5597 USPATFULL

TITLE:

Expression of the developmental I antigen by a cloned

human cDNA encoding a member of a β -1,6-Nacetylglucosaminyltransferase gene family

INVENTOR(S):

Fukuda, Minoru, San Diego, CA, United States

Bierhuizen, Marti F. A., Schiedam, Netherlands

PATENT ASSIGNEE(S):

La Jolla Cancer Research Foundation, La Jolla, CA,

United States (U.S. corporation)

KIND NUMBER DATE -----

PATENT INFORMATION:

US 5484590 19960116 US 1993-118906 19930909 (8)

<--

APPLICATION INFO.:

Utility

DOCUMENT TYPE: FILE SEGMENT:

Granted

PRIMARY EXAMINER: PRIMARY EXAMINER: Wax, Robert A. ASSISTANT EXAMINER: Prouty, Rebecca

LEGAL REPRESENTATIVE: Campbell and Flores

NUMBER OF CLAIMS:

NUMBER OF DRAWINGS:

EXEMPLARY CLAIM:

13 Drawing Figure(s); 11 Drawing Page(s)

LINE COUNT:

1337

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB The present invention provides an isolated nucleic acid molecule encoding both a soluble and membrane-bound human β -1,6-Nacetylglucosaminyltransferase, the I-branching enzyme (IGnT). The invention also provides vectors containing the isolated nucleic acid molecule encoding human IGnT as well as recombinant host cells transformed with the vectors. The invention further provides a method of preparing a membrane-bound form of human IGnT and methods of preparing and purifying soluble human IGnT and active fragments of either form. Also provided are antisense oligonucleotides complementary to a nucleic acid molecule encoding a human IGnT or an active fragment thereof, antibodies directed to the human IGnT, pharmaceutical compositions related to the human IGnT and transgenic nonhuman mammals expressing DNA encoding normal or mutant human IGnT. Also provided are methods for regulating the expression of human IGnT and methods for modifying a biological function mediated by the regulatory activity of human IGnT. Methods of detecting the presence of linear polylactosaminoglycans

expressing i antigenic determinants on a cell surface also are provided.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 12 OF 21 USPATFULL on STN

ACCESSION NUMBER: 94:13514 USPATFULL

TITLE: Inhibitors for replication of retroviruses and for the

expression of oncogene products

INVENTOR(S): Cohen, Jack S., Bethesda, MD, United States

Neckers, Len, Bethesda, MD, United States Stein, Cy, Gaithersburg, MD, United States Loke, She L., Wheaton, MD, United States

Shinozuka, Kazuo, Kazo, Japan

PATENT ASSIGNEE(S): The United States of America as represented by the

Department of Health and Human Services, Washington,

DC, United States (U.S. government)

NUMBER KIND DATE

US 5286717 19940215 US 1992-976777 19921116 (7) PATENT INFORMATION:

APPLICATION INFO.:

RELATED APPLN. INFO.: Division of Ser. No. US 1988-159017, filed on 22 Feb

1988, now abandoned which is a continuation-in-part of

Ser. No. US 1987-30073, filed on 25 Mar 1987, now

abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Rollins, John W.

LEGAL REPRESENTATIVE: Townsend and Townsend Khourie and Crew

NUMBER OF CLAIMS: 20 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 22 Drawing Figure(s); 8 Drawing Page(s)

LINE COUNT: 980

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Phosphorothioate oligodeoxyribonucleotide analogs can be used to prevent

replication of foreign nucleic acids in the presence of normal living cells, as well as to inhibit the proliferation of neoplastic cells.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 13 OF 21 USPATFULL on STN

ACCESSION NUMBER: 94:1412 USPATFULL

TITLE: Inhibitors for replication of retroviruses and for the

expression of oncogene products

INVENTOR(S): Cohen, Jack S., Bethesda, MD, United States

Neckers, Len, Bethesda, MD, United States Stein, Cy, Gaithersburg, MD, United States Loke, Shee L., Wheaton, MD, United States

Shinozuka, Kazuo, Kazo, Japan

PATENT ASSIGNEE(S): The United States of America as represented by the

Department of Health and Human Services, Washington,

DC, United States (U.S. government)

NUMBER KIND DATE ------PATENT INFORMATION:

US 5276019 19940104 US 1988-159017 19880222

APPLICATION INFO.: 19880222 (7)

RELATED APPLN. INFO.: Continuation-in-part of Ser. No. US 1987-30073, filed

on 25 Mar 1987, now abandoned

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Rollins, John W.

LEGAL REPRESENTATIVE: Haight, James C., Ferris, Thomas, Parker, Julie

NUMBER OF CLAIMS: 43 EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 16 Drawing Figure(s); 7 Drawing Page(s)

LINE COUNT: 983

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

Phosphorothioate oligodeoxyribonucleotide analogs can be used to prevent replication of foreign nucleic acids in the presence of normal living cells, as well as to inhibit the proliferation of neoplastic cells.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L15 ANSWER 14 OF 21 USPATFULL on STN

ACCESSION NUMBER:

93:98367 USPATFULL

TITLE:

Inhibitors for replication of retroviruses and for the

expression of oncogene products

INVENTOR(S):

Cohen, Jack S., Bethesda, MD, United States Neckers, Len, Bethesda, MD, United States Stein, Cy, Gaithersburg, MD, United States Loke, She L., Wheaton, MD, United States

Shinozuka, Kazuo, Kazo, Japan

PATENT ASSIGNEE(S):

The United States of America as represented by the Department of Health and Human Services, Washington,

<--

DC, United States (U.S. government)

KIND NUMBER DATE ------

PATENT INFORMATION:

US 5264423 US 1992-976733 19931123

APPLICATION INFO.:

19921116 (7)

Continuation of Ser. No. US 1988-159017, filed on 22 RELATED APPLN. INFO.: Feb 1988, now abandoned which is a continuation-in-part of Ser. No. US 1987-30073, filed on 25 Mar 1987, now

abandoned

DOCUMENT TYPE:

Utility Granted

FILE SEGMENT:

Rollins, John W.

PRIMARY EXAMINER: LEGAL REPRESENTATIVE:

Townsend and Townsend Khourie and Crew

NUMBER OF CLAIMS:

48

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS:

9 Drawing Figure(s); 8 Drawing Page(s)

LINE COUNT:

1018

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB Phosphorothioate oligodeoxyribonucleotide analogs can be used to prevent replication of foreign nucleic acids in the presence of normal living cells, as well as to inhibit the proliferation of neoplastic cells.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

ANSWER 15 OF 21 PCTFULL COPYRIGHT 2005 Univentio on STN

ACCESSION NUMBER:

2001067103 PCTFULL ED 20020822

TITLE (ENGLISH):

FUNCTION HOMOLOGY SCREENING

TITLE (FRENCH):

CRIBLAGE D'HOMOLOGIE DE FONCTIONS

INVENTOR(S):

BERG, Ellen, L.; BUTCHER, Eugene, C.; MELROSE, Jennifer;

PLAVEC, Ivan

PATENT ASSIGNEE(S):

BIOSEEK, INC.; BERG, Ellen, L.; BUTCHER, Eugene, C.; MELROSE, Jennifer;

PLAVEC, Ivan

DOCUMENT TYPE:

Patent

PATENT INFORMATION:

NUMBER KIND DATE

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WO 2001067103 A1 20010913
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DESIGNATED STATES

W:

CF CG CI CM GA GN GW ML MR NE SN TD TG

APPLICATION INFO.: PRIORITY INFO.:

WO 2001-US7190 A 20010306
US 2000-60/186,976 20000306
US 2000-60/195,672 20000407

ABEN A method of screening biologically active agent based on the analysis of complex biological responses in culture. Methods for selecting cells and culture conditions for such screens are provided, as well as the identification of an optimized set of discrete parameters to be measured, and the use of biomap analysis for rapid identification and characterization of drug candidates, genetic sequences acting pathways, and the like. A feature of the invention is simultaneous screening of a large number of cellular pathways, and the rapid identification of

L'invention a trait a une methode de criblage d'un agent biologiquement actif basee sur l'analyse de reponses biologiques complexes en culture. L'invention concerne egalement des methodes de selection de cellules et de conditions de culture pour ces cribles, de meme que l'identification d'un ensemble optimise de parametres distincts a mesurer, et l'utilisation d'une analyse d'une biocarte permettant l'identification et la caracterisation rapides de candidats medicaments, de voies agissant sur les sequences genetiques, et analogue. Une caracteristique de l'invention est le criblage simultane d'un grand nombre de voies cellulaires, ainsi que l'identification rapide de composes provoquant des reponses cellulaires.

L15 ANSWER 16 OF 21 PCTFULL COPYRIGHT 2005 Univentio on STN

ACCESSION NUMBER: TITLE (ENGLISH):

2000075356 PCTFULL ED 20020515 RNA POLYMERASE CHAIN REACTION

TITLE (FRENCH):

AMPLIFICATION EN CHAINE D'ARN PAR POLYMERASE

INVENTOR(S):

LIN, Shi-Lung; YING, Shao-Yao;

CHUONG, Cheng-Ming;

WIDELITZ, Randall, BruceRP : CHAN, Raymond

PATENT ASSIGNEE(S):

LIN, Shi-Lung; YING, Shao-Yao; CHUONG, Cheng-Ming; WIDELITZ, Randall, Bruce

LANGUAGE OF PUBL.: DOCUMENT TYPE:

English Patent

compounds that cause cellular responses.

PATENT INFORMATION:

NUMBER KIND DATE

WO 2000075356 A1 20001214

DESIGNATED STATES

W :

AL AM AT AU AZ BA BB BG BR BY CA CH CN CU CZ DE DK EE ES FI GB GD GE GH GM HR HU ID IL IS JP KE KG KP KR KZ LC LK LR LS LT LU LV MD MG MK MN MW MX NO NZ PL PT RO RU SD SE SG SI SK SL TJ TM TR TT UA UG US UZ VN YU ZW GH GM KE LS MW SD SL SZ UG ZW AM AZ BY KG KZ MD RU TJ TM AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA GN GW ML MR NE SN TD TG

DE DI DO CI CO CI CHI GA GN GW MIL MR

APPLICATION INFO.: WO 1999-US12461 A 19990604

ABEN The present invention provides a fast, simple and specific method for

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generating amplified
messenger RNAs from limited messenger RNAs. The principle of this
RNA-polymerase chain reaction
method relies upon the cycling steps of reverse transcription,
denaturation, double-stranded cDNA
synthesis and then in vitro transcription to bring up the amount of
messenger RNAs to two thousand
folds within one round of above procedure. This method is primarily
designed for differential
screening of tissue-specific gene expressions in cell level, cloning
full-length sequences of
unknown gene transcripts, generating probes for hybridization assays,
synthesizing peptides in
vitro, and preparing representative cDNAs for modern gene chip
technology. In conjunction with a
cell fixation and permeabilisation step, a complete full-length cDNA
library can be directly
generated from few single cells without mRNA degradation.
La presente invention concerne une methode rapide, simple et specifique
de production d'ARN
messagers amplifies a partir d'ARN messagers limites. Le principe de
cette methode d'amplification
en chaine d'ARN par polymerase se base sur les etapes cycliques de la
transcription inverse,
denaturation, synthese d'ADNc bicatenaire et ensuite la transcription in
vitro afin d'elever la
quantite d'ARN messagers de 2000 fois en un cycle de la procedure
precitee. Cette methode est concue
essentiellement pour le criblage differentiel d'expressions de genes a
specificite tissulaire au
niveau cellulaire, le clonage de sequence de longueur totale de
transcrits de gene inconnu, la
production de sondes destinees a des dosages d'hybridation, la synthese
de peptides in vitro et la
preparation d'ADNc representatif pour la technologie moderne des puces a
ADN. Simultanement a une
etape de fixation et de permeabilisation cellulaire, une banque d'ADNc
de longueur totale complete
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ANSWER 17 OF 21 COPYRIGHT 2005 Univentio on STN PCTFULL ACCESSION NUMBER:

l'ARNm.

1999027135 PCTFULL ED 20020515

peut etre generee directement a partir de quelques cellules

TITLE (ENGLISH):

METHOD FOR IDENTIFYING AND INHIBITING FUNCTIONAL

individuelles sans degradation de

NUCLEIC ACID MOLECULES IN CELLS

TITLE (FRENCH):

PROCEDE D'IDENTIFICATION ET D'INHIBITION DE MOLECULES

FONCTIONNELLES D'ACIDE NUCLEIQUE DANS DES CELLULES

INVENTOR(S):

ABFR

NILSEN, Timothy, W.; ROBERTSON, Hugh, D.; KINDT, Thomas, J.

PATENT ASSIGNEE(S):

INNOVIR LABORATORIES, INC.

LANGUAGE OF PUBL.:

English

DOCUMENT TYPE:

Patent

PATENT INFORMATION:

NUMBER

KIND DATE

WO 9927135 A2 19990603

DESIGNATED STATES

W:

AU CA JP AT BE CH CY DE DK ES FI FR GB GR IE IT LU MC

NL PT SE

APPLICATION INFO.: PRIORITY INFO.:

WO 1998-US24854

A 19981120

US 1997-976,220

19971121

US 1998-60/079,851 19980330 Two methodologies are provided: the first provides a means for rapidly ABEN and efficiently identifying essential and functional genes; and the second provides a means for obtaining biologically active nucleic molecules (ribozymes, EGSs, and antisense) which can be used to inactivate functional genes. In the first method, a library of EGSs is prepared based on all possible known compositions. In a preferred embodiment, the EGSs are twelve or thirteen-mers for targeting bacterial RNAse to cleave a substrate. This library is added to the cells containing the genes to be screened, for example, i(E. coli). Those cells in which the EGS causes a loss of viability, or other phenotype, are identified. The EGS(s) responsible for the loss of viability are analyzed, and the resulting sequence information used to identify the gene within the known genomic sequences. In the second method, nucleotide molecules with optimal biological activity, for example, directing cleavage of a gene of interest by RNase P, are rapidly identified through the use of a vector including two reporter genes, the first in phase with the gene of interest, and the second as a control to verify that the vector is present in a cell or to aid in selection of cells containing the vector. Those cells where the gene of interest is cleaved by the functional oligonucleotide molecule can then be identified by reference to reporter gene 1. The responsible functional oligonucleotide molecules is then isolated and characterized. These methods provide powerful tools for identifying essential genes whose sequence is known only as part of a genome with unknown function, as well as means for identifying functional oligonucleotide molecules, useful as diagnostic reagents and therapeutics. ABFR L'invention concerne deux methodologies. Dans la premiere, un moyen pour identifier rapidement et efficacement des genes fonctionnels et essentiels est prevu. Dans la deuxieme, un moyen pour produire des molecules nucleiques biologiquement actives (ribozymes, sequences guides externes et antisens) qui peuvent etre utilisees pour l'inactivation de genes fonctionnels. Dans le premier procede, une banque de sequences guides externes (EGS) a base de toutes les compositions possibles connues est preparee. Dans un mode de realisation prefere, les EGS constituent douze ou treize meres pour cibler l'RNase bacterien pour le clivage d'un substrat. Cette banque est ajoutee a des cellules contenant les genes a cribler, par exemple, i(E. coli). Les cellules dans lesquelles EGS provoque une perte de viabilite, ou d'autre phenotype, sont identifiees. La ou les EGS responsables de la perte de viabilite sont analysees, et l'information relative a la

exemple, dirigeant le clivage d'un gene particulier par RNase P, sont rapidement identifiees au moyen

pour l'identification du gene dans des sequences genomiques connues.

molecules nucleotidiques ayant une activite biologique optimale, par

sequence resultante est utilisee

Dans le deuxieme procede, les

d'un vecteur comprenant deux

genes reporters, le premier en phase avec le gene en question et le deuxieme permettant de verifier

que le vecteur est present dans une cellule ou facilitant la selection de cellules contenant le

vecteurs. Les cellules dans lesquelles le gene en question est clive par la molecule

oligonucleotidique fonctionnelle peuvent ensuite etre identifiees en fonction du gene reporter 1.

Les molecules oligonucleotidiques fonctionnelles responsables sont ensuite isolees et caracterisees.

Lesdits procedes constituent des outils puissants pour l'identification de genes essentiels dont la

sequence est connue seulement comme faisant partie integrante d'un genome a fonction inconnue, ainsi

que des moyens d'identification de molecules oligonucleotidiques fonctionnelles, utiles comme

reactifs diagnostiques et comme agents therapeutiques.

L15 ANSWER 18 OF 21 PCTFULL COPYRIGHT 2005 Univentio on STN

ACCESSION NUMBER: 1996018733 PCTFULL ED 20020514

TITLE (ENGLISH): RIBOZYME-MEDIATED INACTIVATION OF LEUKEMIA-ASSOCIATED

RNA

TITLE (FRENCH): INACTIVATION INDUITE PAR RIBOZYME DE L'ARN ASSOCIE A LA

LEUCEMIE

INVENTOR(S): PACE, Umberto;

GEORGE, Shaji, T.; GOLDBERG, Allan, R.

PATENT ASSIGNEE(S): INNOVIR LABORATORIES, INC.

LANGUAGE OF PUBL.: English DOCUMENT TYPE: Patent

PATENT INFORMATION:

NUMBER KIND DATE

WO 9618733 A2 19960620

DESIGNATED STATES

W: AU CA JP AT BE CH DE DK ES FR GB GR IE IT LU MC NL PT

SE

APPLICATION INFO.: WO 1995-US16451 A 19951214 PRIORITY INFO.: US 1994-354,956 19941214

ABEN RNA molecules, including ribozymes, external guide sequences for RNAse P, and antisense

oligonucleotides have been constructed which promote ribozyme cleavage of, or block transcription $% \left(1\right) =\left(1\right) \left(1$

of, respectively, specific cancer-associated RNA, for example, acute promyeloleukocytic

leukemia-associated RNA, follicular lymphoma-associated RNA and chronic myelocytic

leukemia-associated RNA. Methods of producing and using such RNA molecules are also described.

ABFR La presente invention concerne la construction de molecules d'ARN, y compris de ribozymes, de

sequences guides externes pour la ribonuclease ${\tt P}$ et d'oligonucleotides antisens, lesquelles

molecules, suivant le cas, favorisent la coupure par ribozymes d'ARN specifiques de cancers, ou en

bloquent la transcription. Ces ARN sont notamment l'ARN de la leucemie promyeloleucocytaire aique,

l'ARN du lymphome folliculaire, et l'ARN de la leucemie myeloloide chronique. L'invention concerne

egalement des procedes de production et d'utilisation de telles molecules d'ARN.

L15 ANSWER 19 OF 21 PCTFULL COPYRIGHT 2005 Univentio on STN

ACCESSION NUMBER: 1995007020 PCTFULL ED 20020514

TITLE (ENGLISH): EXPRESSION OF THE DEVELOPMENTAL I ANTIGEN BY A CLONED

HUMAN CDNA ENCODING A BETA-1,6-N-

ACETYLGLUCOSAMINYLTRANSFERASE

TITLE (FRENCH): EXPRESSION DE L'ANTIGENE I DE DEVELOPPEMENT PAR UN ADNC

CODANT UNE BETA-1,6-N-ACETYLGLUCOSAMINYLTRANFERASE

INVENTOR(S): FUKUDA, Minoru;

BIERHUIZEN, Marti, F., A.

PATENT ASSIGNEE(S): LA JOLLA CANCER RESEARCH FOUNDATION

LANGUAGE OF PUBL: English

DOCUMENT TYPE: Engils

PATENT INFORMATION:

NUMBER KIND DATE

WO 9507020 A1 19950316

DESIGNATED STATES

W: CA JP

APPLICATION INFO.: WO 1993-US8476 A 19930909

ABEN The present invention provides an isolated nucleic acid molecule

encoding both a soluble and

membrane-bound human 'beta'-1,6-N-acetylglucosaminyltransferase, the

I-branching enzyme (IGnT). The

invention also provides vectors containing the isolated nucleic acid

molecule encoding human IGnT as

well as recombinant host cells transformed with the vectors. The

invention further provides a method

of preparing a membrane-bound form of human IGnT and methods of preparing and purifying soluble

human IGnT and active fragments of either form. Also provided are antisense oligonucleotides

complementary to a nucleic acid molecule encoding a human IGnT or an active fragment thereof,

antibodies directed to the human IGnT, pharmaceutical compositions related to the human IGnT and

transgenic nonhuman mammals expressing DNA encoding normal or mutant human IGnT. Also provided are

methods for regulating the expression of human IGnT and methods for modifying a biological function

mediated by the regulatory activity of human IGnT. Methods of detecting the presence of linear

polylactosaminoglycans expressing i antigenic determinants on a cell surface also are provided.

ABFR La presente invention concerne une molecule d'acide nucleique codant a la fois les formes

solubles et les formes liees a l'enveloppe de

l'acetylglucosaminyltransferase, l'enzyme ramifiante-I

(IGnT). On decrit egalement des vecteurs contenant la molecule d'acide nucleique isolee codant

l'IGnT humaine ainsi que les cellules hotes de recombinaison transformees a l'aide de vecteurs. De

plus, l'invention se rapporte a un procede de preparation d'une forme liee de l'IGnT humaine et a

des procedes de preparation et de purification de l'IGnT humaine soluble et des fragments actifs de

chaque forme. On decrit egalement des oligonucleotides anti-sens complementaires a une molecule

d'acide nucleique codant une IGnT humaine ou bien un fragment actif de celle-ci, des anticorps

diriges contre l'IGnT humaine, des compositions pharmaceutiques apparentees a l'IGnT humaine et des

mammiferes non humains transgeniques exprimant de l'ADN codant une IGnT humaine normale ou mutante.

On decrit encore des procedes de regulation de l'expression de l'IGnT et des procedes de

modification d'une fonction biologique induite par l'activite regulatrice de l'IGnT. De plus, on

decrit des procedes de detection de la presence de

polylactosaminoglycanes exprimant des determinants antigeniques sur la surface d'une cellule.

ANSWER 20 OF 21 PCTFULL

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ACCESSION NUMBER:

1995006062 PCTFULL ED 20020514

TITLE (ENGLISH):

NOVEL PROTEINS ISOLATED FROM NERVE CELLS, DNA SEQUENCES

ENCODING SAME AND USAGES THEREOF

TITLE (FRENCH):

NOUVELLES PROTEINES ISOLEES A PARTIR DE CELLULES NERVEUSES, SEQUENCES D'ADN CODANT CES PROTEINES ET

LEURS UTILISATIONS

INVENTOR(S):

LIN, Siang-Yo;

WU, Kuo;

BLACK, Ira, B.

PATENT ASSIGNEE(S):

THE UNIVERSITY OF MEDICINE AND DENTISTRY OF NEW JERSEY

PATENT INFORMATION:

NUMBER KIND DATE

WO 9506062

A1 19950302

DESIGNATED STATES

W :

DOCUMENT TYPE:

AT BE CH DE DK ES FR GB GR IE IT LU MC NL PT SE

APPLICATION INFO.: PRIORITY INFO.:

WO 1994-US9601 A 19940823 US 1993-8/110,501 19930823 US 1994-8/188,422 19940124 US 1994-8/276,357 19940718

ABEN The present invention is based on the identification of four novel protein tyrosine kinases

(PTKs) isolated from the postsynaptic density (PSD). These PTKs, whose molecular weights were

approximately 166, 90, 66, and 50 kDa, as shown in the figure, were found to be resistant to

Genistein, did not crossreact with antibodies which bind to other PTKs known in the art. The present

invention further discloses previously unidentified proteins, isolated from the PSD, which have

molecular weights of approximately 110, 120 and 270 kDa. These proteins were found to crossreact

with an antibody which is selective for dystrophin.

ABFR La presente invention se fonde sur l'identification de quatre nouvelles proteines-tyrosine-kinases (PTK) isolees a partir de la densite postsynaptique (DPS). On a decouvert

que ces PTK dont les masses moleculaires sont de 166, 90, 66 et 50 kDa approximativement comme

represente dans la figure, sont resistantes au Genistein, et ne presentent pas de reaction croisee

avec les anticorps qui se lient a d'autres PTK connues dans la technique actuelle. De plus, la

presente invention concerne des proteines non identifiees auparavant, isolees a partir de la DPS,

presentant des masses moleculaires de 110, 120 et 270 kDa approximativement. On a trouve que ces

proteines presentaient une reaction croisee avec un anticorps qui est selectif pour la dystrophine.

L15 ANSWER 21 OF 21

PCTFULL COPYRIGHT 2005 Univentio on STN

ACCESSION NUMBER:

1994024284 PCTFULL ED 20020513

TITLE (ENGLISH):

HUMAN N-METHYL-D-ASPARTATE RECEPTOR SUBUNITS, NUCLEIC

ACIDS ENCODING SAME AND USES THEREFOR

SOUS-UNITES RECEPTRICES DU N-METHYL-D-ASPARTATE HUMAIN, TITLE (FRENCH):

ACIDES NUCLEIQUES CODANT POUR ELLES, ET LEUR

UTILISATION

INVENTOR(S): DAGGETT, Lorrie, P.;

ELLIS, Steven, B.; LIAW, Chen, Wang;

LU, Chin-Chun

THE SALK INSTITUTE BIOTECHNOLOGY/INDUSTRIAL ASSOCIATES, PATENT ASSIGNEE(S):

INC.;

DAGGETT, Lorrie, P.; ELLIS, Steven, B.; LIAW, Chen, Wang; LU, Chin-Chun

LANGUAGE OF PUBL.:

DOCUMENT TYPE:

English Patent

PATENT INFORMATION:

NUMBER KIND DATE

WO 9424284 A1 19941027

DESIGNATED STATES

W:

AT AU BB BG BR BY CA CH CN CZ DE DK ES FI GB GE HU JP KG KP KR KZ LK LU LV MD MG MN MW NL NO NZ PL PT RO RU SD SE SI SK TJ TT UA US UZ VN AT BE CH DE DK ES FR GB GR IE IT LU MC NL PT SE BF BJ CF CG CI CM GA GN ML MR NE SN TD TG

APPLICATION INFO.:

PRIORITY INFO.:

WO 1994-US4387 A 19940420 US 1993-8/052,449 19930420

ABEN In accordance with the present invention, there are provided nucleic acids encoding human NMDA

receptor protein subunits and the proteins encoded thereby. The NMDA receptor subunits of the

invention comprise components of NMDA receptors that have cation-selective channels and bind

glutamate and NMDA. In one aspect of the invention, the nucleic acids encode NMDAR1 and NMDAR2

subunits of human NMDA receptors. In a preferred embodiment, the invention nucleic acids encode

NMDAR1, NMDAR2A, NMDAR2B, NMDAR2C and NMDAR2D subunits of human NMDA receptors. In addition to being

useful for the production of NMDA receptor proteins subunit, these nucleic acids are also useful as

probes, thus enabling those skilled in the art, without undue experimentation, to identify and

isolate related human receptor subunits. Functional glutamate receptors can be assembled, in

accordance with the present invention, from a plurality of one type of NMDA receptor subunit protein

(homomeric) or from a mixture of two or more types of subunit proteins (heteromeric). In addition to

disclosing novel NMDA receptor protein subunits, the present invention also comprises methods for

using such receptor subunits to identify and characterize compounds which affect the function of

such receptors, e.g., agonists, antagonists, and modulators of glutamate receptor function. The

invention also comprises methods for determining whether unknown protein(s) are functional as NMDA

receptor subunits. Acides nucleiques codant pour les sous-unites des proteines receptrices ABFR du NMDA humain et

proteines codees par ce moyen. Les sous-unites des recepteurs du NMDA de l'invention comprennent des

composants des recepteurs du NMDA a canaux selecteurs de cations qui

```
fixent le glutamate et le NMDA.
Dans l'une des realisations de l'invention, les acides nucleiques codent
pour les sous-unites NMDAR1
et NMDAR2 des recepteurs du NMDA humain. Dans la realisation preferee,
les acides nucleiques objets
de l'invention codent pour les sous-unites NMDAR1, NMDAR2A, NMDAR2B,
NMDAR2C et NMDAR2D des
recepteurs du NMDA humain. En plus de leur utilite pour la production de
sous-unites des proteines
receptrices du NMDA, lesdits acides nucleiques peuvent egalement servir
de sondes permettant aux
inities d'identifier et d'isoler, sans devoir recourir a une
experimentation superflue, les
sous-unites associees des recepteurs humains. Selon la presente
invention, les recepteurs
fonctionnels du glutamate peuvent etre assembles a partir d'une
multiplicite d'un type de
sous-unites de proteine receptrice du NMDA (homomeres) ou a partir d'un
melange d'une ou plusieurs
sous-unites de proteines (heteromeres). En plus de la presentation de
nouvelles sous-unites de la
proteine receptrice du NMDA, la presente invention comporte des prodedes
d'utilisation des
sous-unites de recepteurs pour identifier et caracteriser des composes
influant sur le
fonctionnement de ces recepteurs, par exemple des agonistes, des
antagonistes ou des modulateurs de
la fonction des recepteurs du glutamate. L'invention comporte egalement
des methodes pour determiner
si des proteines inconnues ont la fonction de sous-unites de recepteurs
de NMDA.
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=> d his

(FILE 'HOME' ENTERED AT 13:37:56 ON 16 JUN 2005)

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FILE 'MEDLINE, BIOSIS, SCISEARCH, EMBASE, CA, USPATFULL, PCTFULL' ENTERED
     AT 13:39:40 ON 16 JUN 2005
L1
          31952 S ((MRNA OR RNA OR VRNA) (N) (ADNA OR CDNA OR DNA)) (3N) HYBRID
L2
       10362864 S L1 (S) INHIB? OR REDUC?
L3
           3102 S L1 (S) (INHIB? OR REDUC?)
L4
             67 S L1 (S) ((INHIB? OR REDUC?) (N) EXPRES?)
L5
             67 DUP REM L4 (0 DUPLICATES REMOVED)
L6
             14 S (L5 AND PY<=2001)
L7
          35458 S LIN, S?/AU
L8
            797 S CHUONG, C?/AU
            216 S WIDELITZ, R?/AU
Ь9
L10
          36265 S L7 OR L8 OR L9
L11
             26 S L10 AND L1
L12
            18 DUP REM L11 (8 DUPLICATES REMOVED)
L13
             32 S L12 OR L6
L14
            32 DUP REM L13 (0 DUPLICATES REMOVED)
L15
            21 S L14 AND PY<=2001
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